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10 **UNITED STATES DISTRICT COURT**
11 **NORTHERN DISTRICT OF CALIFORNIA**
12 **OAKLAND DIVISION**

13 STEVEN STREZSAK, Individually and on
14 Behalf of All Others Similarly Situated,

15 Plaintiff,

16 v.

17 ARDELYX INC., MIKE RAAB, and JUSTIN
18 RENZ,

19 Defendants.

Case No. _____

CLASS ACTION COMPLAINT

JURY TRIAL DEMANDED

Plaintiff Steven Strezsak (“Plaintiff”) makes the following allegations, individually and on behalf of all other similarly situated, by and through Plaintiff’s counsel, upon information and belief, except as to those allegations concerning Plaintiff, which are alleged upon personal knowledge. Plaintiff’s information and belief is based upon, *inter alia*, counsel’s investigation, which included, among other things, review and analysis of: (i) regulatory filings made by Ardelyx Inc. (“Ardelyx” or the “Company”) with the SEC; (ii) press releases and media reports issued by and disseminated by the Company; and (iii) analyst reports, media reports, and other publicly disclosed reports and information about the Company. Plaintiff believes that substantial evidentiary support will exist for the allegations set forth herein, after a reasonable opportunity for discovery.

SUMMARY OF THE ACTION

1. Plaintiff brings this federal securities class action under §§10(b) and 20(a) of the Securities Exchange Act of 1934 (the “Exchange Act”) and SEC Rule 10b-5 promulgated thereunder, 17 C.F.R. §240.10b-5, on behalf of a class consisting of all persons and entities, other than Defendants and their affiliates, who purchased Ardelyx securities between August 6, 2020 and July 19, 2021, inclusive (the “Class Period”), and who were damaged thereby (the “Class”).

2. Ardelyx is a specialized biopharmaceutical company focused on developing first-in-class medicine to improve treatment for people with cardiorenal disease. This includes patients with chronic kidney disease (“CKD”) on dialysis suffering from elevated serum phosphorus, or hyperphosphatemia; and CKD patients and/or heart failure patients with elevated serum potassium, or hyperkalemia.

3. In June 2020, Defendants submitted a New Drug Application (“NDA”) to the U.S. Food and Drug Administration (“FDA”) for Ardelyx’s lead product candidate, tenapanor, a supposedly first-in-class medicine for the control of serum phosphorus in adult patients with CKD on dialysis. According to Ardelyx, tenapanor has “a unique mechanism of action and acts locally in the gut to inhibit the sodium hydrogen exchanger 3, or NHE3,” resulting in the “tightening of the epithelial cell junctions, thereby significantly reducing paracellular uptake of phosphate, the

1 primary pathway of phosphate absorption.” If approved, tenapanor “would be the first therapy for
2 phosphate management that blocks phosphorus absorption at the primary pathway of uptake[.]”
3 and “could greatly improve patient adherence and compliance with one single pill dosed twice
4 daily in contrast to current therapies where typically multiple pills are taken before every meal.”
5 Thus, tenapanor (and its promise) was widely touted by Defendants and, accordingly, extremely
6 important to the valuation (and future success) of Ardelyx securities.

7 4. The FDA accepted Ardelyx’s NDA in September 2020 and set a Prescription Drug
8 User Fee Act (“PDUFA”) date of April 29, 2021.

9 5. The Company repeatedly lauded this development, highlighting the FDA’s
10 acceptance and review of the NDA, supported by so-called “successful” Phase 3 studies, in each
11 subsequently filed quarterly report and in the Company’s 2020 Annual Report (defined below).
12 Even when the FDA requested that the Company provide additional information related to
13 Ardelyx’s clinical data, which caused the PDUFA date to slip by three months, Defendants
14 continued to hype Ardelyx’s “positive” clinical trial results, which, according to them, showed
15 “improvements” over current treatments, supported tenapanor’s “clinical safety and efficacy,” and
16 reinforced its “potential” as a “transformative” treatment. At no point did Defendants state (much
17 less suggest) that there may be fatal issues with the drug, its clinical trial data, or both. Rather,
18 Defendants simply claimed that the FDA’s request was merely because they needed help to “better
19 understand the clinical data in light of tenapanor’s novel mechanism of action as compared to
20 approved therapies.”

21 6. Defendants’ rosy narrative, however, came to a screeching halt after the market
22 closed on July 19, 2021. At that time, Ardelyx announced that it had received a letter from the
23 FDA, dated July 13, 2021, that said the administration had found deficiencies that precluded
24 discussion around the would-be labeling and post-marketing requirements for tenapanor.
25 Critically, the FDA said it *detected issues with both the size and clinical relevance* of the drug’s
26 treatment effect.

7. Immediately, analysts cut their price targets and downgraded the Company's rating. Piper Sandler, for example, rated Ardelyx neutral (down from a buy-equivalent rating) and wrote, "we struggle to see a path forward for Tenapanor." Raymond James, another analyst, reset the Company's price target to \$4 from \$14 per share.

8. The Company's share price likewise plunged, falling \$9.71 per share, or nearly 74%, in a single day, to close at \$2.01 per share on July 20, 2021, before falling another 4.2% by market close on July 21, 2021.

9. Throughout the Class Period, Defendants made materially false and misleading statements regarding tenapanor and the likelihood that it would be approved by the FDA. Defendants possessed, were in control over, and, as a result, knew (or had reason to know) that the data submitted to support the NDA was insufficient in that it showed a lack of clinical relevance of the drug's treatment effect, making it foreseeably likely (if not certain) that the FDA would not approve the drug.

10. This lawsuit seeks to recover damages sustained as a result of Defendants' wrongdoing.

JURISDICTION AND VENUE

11. The claims asserted herein arise under §§10(b) and 20(a) of the Exchange Act, 15 U.S.C. §§78j(b) and 78t(a), and SEC Rule 10b-5 promulgated thereunder, 17 C.F.R. §240.10b-5.

12. This Court has jurisdiction over the subject matter of this action pursuant to 28 U.S.C. §§1331 and §27 of the Exchange Act, 15 U.S.C. §78aa.

13. The Court has jurisdiction over each of the Defendants named herein because each is an individual or a corporation who has sufficient minimum contacts with this District so as to render the exercise of jurisdiction by the District Court permissible under traditional notions of fair play and substantial justice.

14. Venue is proper in this District pursuant to §27 of the Exchange Act, 15 U.S.C. §78aa and 28 U.S.C. §1391(b), as the Company's headquarters are located within this District.

Defendants knew that the adverse facts specified herein had not been disclosed to and were being concealed from the public and that the positive representations being made were false and misleading. The Individual Defendants are liable for the false statements pleaded herein.

SUBSTANTIVE ALLEGATIONS

22. Ardelyx is a biotechnology company focused on the development of therapies for cardiorenal disorder. Though Ardelyx's lead product candidate, tenapanor, has been approved by the FDA as a treatment for irritable bowel syndrome associated constipation, the Company has not commercialized it in the United States nor generated any revenue from its sale. Rather, Ardelyx has focused on advancing another indication for the drug, namely for helping to control serum phosphorus in adult CKD patients on dialysis.

23. In fact, Ardelyx presented tenapanor to the FDA as a new treatment to manage hyperphosphatemia in CKD patients undergoing dialysis treatment based on a Phase 3 program for the control of serum phosphorus in CKD patients on dialysis. In December 2019, the Company reported (purportedly) statistically significant topline efficacy results from its second monotherapy Phase 3 clinical trial, the PHREEDOM trial, which had followed a "successful" monotherapy Phase 3 clinical trial completed in 2017 that (again, purportedly) achieved statistical significance for the primary endpoint.¹

24. Consequently, obtaining regulatory approvals for tenapanor for the control of serum phosphorus in adult CKD patients on dialysis was critical.

MATERIALLY FALSE AND MISLEADING STATEMENTS

25. The Class Period begins on August 6, 2020, when Ardelyx issued a press release announcing that it submitted an NDA to the FDA for the review of tenapanor as a first-in-class therapy to control serum phosphorus in adult patients with CKD on dialysis. According to the

¹ PHREEDOM was a one-year study with a 26-week *open-label* treatment period and a 12-week double-blind, placebo-controlled randomized withdrawal period followed by a 14-week *open-label* safety extension period. An active safety control group, for safety analysis only, received sevelamer, *open-label*, for the entire 52-week study period. Patients completing the PHREEDOM trial from both the tenapanor arm and the sevelamer active safety control arm had the option to participate in NORMALIZE, an ongoing *open-label* 18-month extension study.

1 press release, the filing was supported by three *successful* Phase 3 studies demonstrating
 2 tenapanor's ability to *reduce* phosphate levels. In addition, the release noted that "additional
 3 *positive data* from the ongoing NORMALIZE Phase 4 study" showed a "58% *improvement* over
 4 current standard of care." [Emphasis added.]

5 26. Also on August 6, 2020, Ardelyx filed with the SEC its quarterly report on Form
 6 10-Q for the period ending June 30, 2020 (the "2Q20 10-Q"), further touting the apparent benefits
 7 of tenapanor, stating in relevant part:

8 In June 2020, we announced *positive* results from a planned interim data analysis
 9 from our ongoing NORMALIZE Phase 4 study evaluating tenapanor, as
 10 monotherapy or in combination with sevelamer, to achieve serum phosphorus
 11 levels in the normal range (2.5 – 4.5 mg/dL) in patients with CKD on dialysis. The
 12 NORMALIZE extension study allowed patients from our PHREEDOM study to
 13 continue therapy with tenapanor and enabled those patients in the PHREEDOM
 14 safety control arm receiving sevelamer carbonate to transition to tenapanor. *The*
 15 *data from the planned interim analysis demonstrated that the foundational use*
 16 *of tenapanor as monotherapy or in combination with sevelamer carbonate*
 17 *produces a significant phosphorus-lowering effect* with a mean serum
 18 phosphorous reduction of 2.33 mg/dL, from a mean baseline phosphorus of 7.27
 19 mg/dL at the beginning of the PHREEDOM trial to a mean of 4.94 mg/dL at the
 20 time of this analysis. Of the 171 patients in this interim analysis who completed up
 21 to 9 months of treatment in this extension study, up to 47.4% achieved a normal
 22 serum phosphorus level, and of those, the majority were on tenapanor alone or
 23 tenapanor with low dose sevelamer of ≤ 3 sevelamer tablets per day. These data
 24 represent a 58% *improvement* in the rate of patients who achieve a normal serum
 25 phosphorus level, as compared to current treatment practice data as reported in the
 26 April 2020 Dialysis Outcomes Practice Patterns Study ("DOPPS") Practice
 27 Monitor.

18 * * *

19 Tenapanor, if approved, would be the first therapy for phosphate management that
 20 blocks phosphorus absorption at the primary pathway of uptake. It is not a
 21 phosphate binder. *Tenapanor is a novel, potent, small molecule, that has been*
 22 *shown in the phase 3 studies to treat hyperphosphatemia* as monotherapy and as
 23 a dual mechanism approach. Additionally, as such we believe tenapanor could
 24 greatly improve patient adherence and compliance with one single pill dosed twice
 25 daily in contrast to current therapies where typically multiple pills are taken before
 26 every meal.

27 [Emphasis added].

28 27. On November 5, 2020, Ardelyx filed with the SEC on Form 10-Q its third quarter
 2020 financial results, substantially repeating the same claims made in the Company's 2Q20 10-
 Q. Defendants also issued a press release that emphasized certain "business highlights," including

that the FDA accepted the NDA submitted by Defendants for tenapanor to control serum phosphorus in adult patients with CKD on dialysis. Defendants, again, claimed that the filing was supported by three *successful* Phase 3 studies *demonstrating tenapanor's ability to reduce* phosphate levels, with Defendant Raab, specifically, touting "clinical data presented at ASN Kidney Week 2020[, which] *support[s] the clinical safety and efficacy of tenapanor and reinforce[s] its potential* to transform the treatment landscape for patients." [Emphasis added.]

28. On March 8, 2021, Ardelyx filed with the SEC on Form 10-K its Fourth Quarter and Full year 2020 Financial Results, which touted the Company's ability to monetize tenapanor upon FDA approval. For example, it stated:

Tenapanor: A New Approach for The Control of Serum Phosphorus in CKD Patients on Dialysis

Our portfolio is led by the development of tenapanor, a first-in-class medicine for the control of serum phosphorus in adult patients with CKD on dialysis. Tenapanor for the control of serum phosphorus has a unique mechanism of action and acts locally in the gut to inhibit the sodium hydrogen exchanger 3 ("NHE3"). This results in the tightening of the epithelial cell junctions, thereby significantly reducing paracellular uptake of phosphate, the primary pathway of phosphate absorption. On September 15, 2020 we announced that the FDA accepted the filing of our NDA for tenapanor for the control of serum phosphorus in adult patients with CKD on dialysis. The acceptance of our NDA represents the next critical step toward *bringing to market* a completely new approach to the management of hyperphosphatemia. The FDA has set a PDUFA date of April 29, 2021. *We continue to advance commercial preparations for the launch of tenapanor for this indication.* The NDA is supported by three *successful* Phase 3 trials involving over 1,000 patients that evaluated the use of tenapanor for the control of serum phosphorus in CKD patients on dialysis, with two trials evaluating tenapanor as monotherapy and one trial evaluating tenapanor as part of a dual mechanism approach with phosphate binders.

* * *

In December 2019, we reported *statistically significant* topline efficacy results from our second monotherapy Phase 3 clinical trial, the PHREEDOM trial, which evaluated tenapanor for the control of serum phosphorus in CKD patients on dialysis. The PHREEDOM trial followed a *successful* monotherapy Phase 3 clinical trial completed in 2017, the BLOCK trial, which achieved *statistical significance* for the primary endpoint. The only adverse event reported in these Phase 3 trials in greater than 5% of patients was diarrhea, with an incidence rate of 52% in the PHREEDOM trial and 39% in the BLOCK trial, with most incidences in each trial being mild to moderate in nature. PHREEDOM is a one-year study with a 26-week *open-label* treatment period and a 12-week double-blind, placebo-controlled randomized withdrawal period followed by a 14-week *open-label* safety extension period. An active safety control group, for safety analysis only, received sevelamer, open-label, for the entire 52-week study period. Patients completing the PHREEDOM trial from both the tenapanor arm and the sevelamer active safety

1 control arm had the option to participate in NORMALIZE, an ongoing *open-label*
2 18-month extension study.

3 In June 2020, we announced **positive** results from a planned analysis from our
4 ongoing NORMALIZE extension study evaluating tenapanor, as monotherapy or
5 in combination with sevelamer, to achieve serum phosphorus levels in the normal
6 range (2.5 – 4.5 mg/dL) in patients with CKD on dialysis. The NORMALIZE
7 extension study allowed patients from our PHREEDOM study to continue therapy
8 with tenapanor and enabled those patients in the PHREEDOM safety control arm
receiving sevelamer carbonate to transition to tenapanor. The data from the planned
interim analysis demonstrated that the foundational use of tenapanor as
monotherapy or in combination with sevelamer carbonate produces a significant
phosphorus-lowering effect with a mean serum phosphorous reduction of 2.33
mg/dL, from a mean baseline phosphorus of 7.27 mg/dL at the beginning of the
PHREEDOM trial to a mean of 4.94 mg/dL at the time of this analysis.

9 [Emphasis added.]

10 29. Also on March 8, 2021, Ardelyx issued a press release within which Defendant
11 Raab stated: “[t]he stage is set for an exciting year for Ardelyx in 2021,” since “***we are well***
12 ***positioned and well prepared to commercialize*** tenapanor upon potential FDA approval of the
13 first and only phosphate absorption inhibitor for the control of serum phosphorus.” [Emphasis
14 added.]

15 30. Then, on April 29, 2021, Ardelyx issued a press release announcing the need to
16 provide additional analyses of its clinical data to the FDA in connection with the FDA’s ongoing
17 review of the Company’s NDA for tenapanor. According to the Company, the FDA requested this
18 information to help it “better understand the clinical data in light of tenapanor’s novel mechanism
19 of action as compared to approved therapies.” Since this information constituted a “major
20 amendment to the NDA,” the PDUFA date was extended three months to July 29, 2021.

21 31. Defendant Raab offered an optimistic take on the FDA’s request in a May 6, 2021
22 press release announcing the Company’s First Quarter 2021 Financial Results, stating in relevant
23 part:

24 We continue to prepare for the potential approval and launch of tenapanor
25 following the recent extension of our PDUFA date to July. ***We remain confident***
26 ***in the comprehensive data included in our New Drug Application*** and believe
27 tenapanor represents an attractive alternative to currently available therapies to
control serum phosphorus in CKD patients on dialysis. To that end, we are
committed to working with the FDA through the completion of its review of our
NDA and ***look forward to the possibility of making a significant impact*** in the
28 lives of patients.

1 [Emphasis added.]

2 32. The statements identified above were materially false and misleading and failed to
3 disclose material facts about tenapanor and the likelihood that it would be approved by the FDA.
4 Defendants possessed, were in control over, and, as a result, knew (or had reason to know) that the
5 data submitted to support the NDA was insufficient in that it showed a lack of clinical relevance
6 of the drug's treatment effect, making it foreseeably likely (if not certain) that the FDA would not
7 approve the drug.

8 **THE TRUTH EMERGES**

9 33. Defendants' upbeat narrative came to a screeching halt after the markets closed on
10 July 19, 2021, when they announced that Ardelyx received a letter from the FDA *on July 13, 2021*,
11 stating that "the FDA *has identified deficiencies that preclude discussion of labeling and post-*
12 *marketing requirements/commitments.*" In particular, the FDA noted that "*a key issue is the size*
13 *of the treatment effect and its clinical relevance.*" [Emphasis added.]

14 34. On this news, the price of Ardelyx's shares plunged from their July 19, 2021 closing
15 price of \$7.70 per share to a July 20, 2021 close of just \$2.01 each. This represents a one-day drop
16 of nearly 74%, or hundreds of millions of dollars in lost market capitalization.

17 **CLASS ACTION ALLEGATIONS**

18 35. Plaintiff repeats and realleges each and every allegation contained above as if fully
19 set forth herein.

20 36. Plaintiff brings this action as a class action, pursuant to Rules 23(a) and 23(b)(3) of
21 the Federal Rules of Civil Procedure, on behalf of the Class, consisting of all persons and entities
22 that purchased, or otherwise acquired, the common stock of Ardelyx during the Class Period.

23 37. Excluded from the Class are: (i) Defendants; (ii) present or former executive
24 officers of Ardelyx, members of the Board, and members of their immediate families (as defined
25 in 17 C.F.R. §229.404, Instructions (1)(a)(iii) and (1)(b)(ii)); (iii) any of the foregoing persons'
26 legal representatives, heirs, successors, or assigns; and (iv) any entities in which Defendants have
27 or had a controlling interest, or any affiliate of Ardelyx.

1 38. The members of the Class are so numerous that joinder of all members is
 2 impracticable. Throughout the Class Period, the Company's common stock was actively traded
 3 on the NASDAQ, a national securities exchange. While the exact number of Class members is
 4 unknown to Plaintiff at this time, and can only be ascertained through appropriate discovery,
 5 Plaintiff believes that there are hundreds or thousands of members in the Class. Millions of
 6 Ardelyx shares were publicly traded during the Class Period on the NASDAQ. Record owners
 7 and other members of the Class may be identified from records maintained by Ardelyx or its
 8 transfer agent and may be notified of the pendency of this action by mail, using a form of notice
 9 similar to that customarily used in securities class actions.

10 39. Plaintiff's claims are typical of the claims of Class members, who were all similarly
 11 affected by Defendants' wrongful conduct in violation of the federal securities laws. Further,
 12 Plaintiff will fairly and adequately protect the interests of Class members and have retained
 13 counsel competent and experienced in class and securities litigation.

14 40. Common questions of law and fact exist as to all members of the Class and
 15 predominate over any questions solely affecting individual members of the Class. Among the
 16 questions of law and fact common to the members of the Class are:

- 17 (a) whether Defendants violated the Exchange Act;
- 18 (b) whether Defendants' statements to the investing public during the Class Period
- 19 omitted and/or misrepresented material facts;
- 20 (c) whether Defendants' statements to the investing public during the Class Period
- 21 omitted material facts necessary in order to make the statements made, in light
- 22 of the circumstances under which they were made, not misleading;
- 23 (d) whether Defendants knew or recklessly disregarded that their statements were
- 24 false and misleading;
- 25 (e) whether the price of Ardelyx's common stock was artificially inflated; and
- 26 (f) the extent of damage sustained by Class members and the appropriate measure
- 27 of damages.

41. A class action is superior to all other available methods for the fair and efficient adjudication of this controversy, since joinder of all members is impracticable. Further, as the damages suffered by individual Class members may be relatively small, the expense and burden of individual litigation makes it impossible for Class members to individually redress the wrongs done to them. There will be no difficulty in the management of this action as a class action.

**For Violations of §10(b) of the Exchange Act and
Rule 10b-5 Promulgated Thereunder
(Against all Defendants)**

42. Plaintiff repeats and realleges each and every allegation contained above as if fully set forth herein.

43. This Count is asserted on behalf of all members of the Class against Ardelyx and the Individual Defendants for violations of §10(b) of the Exchange Act, 15 U.S.C. §78(b), and Rule 10b-5 promulgated thereunder, 17 C.F.R. §240.10b-5.

44. These Defendants carried out a plan, scheme, and course of conduct which was intended to, and did: (i) deceive the investing public, including Plaintiff and the other Class members, as alleged herein; and (ii) caused Plaintiff and the other members of the Class to purchase Ardelyx securities at artificially inflated prices. In furtherance of this unlawful scheme, plan, and course of conduct, each of these Defendants took the actions set forth herein.

45. During the Class Period, Defendants disseminated or approved the false statements specified herein, among others, which they knew, or deliberately disregarded, were materially misleading in that they contained material misrepresentations and failed to disclose material facts necessary in order to make the statements made, in light of the circumstances under which they were made, not misleading.

46. Defendants: (i) employed devices, schemes, and artifices to defraud; (ii) made untrue statements of material fact and/or omitted to state material facts necessary to make the statements made not misleading; and (iii) engaged in acts, practices, and a course of business that operated as a fraud and deceit upon the purchasers of the Company's securities in an effort to

1 maintain artificially high market prices for Ardelyx securities in violation of §10(b) of the
2 Exchange Act and Rule 10b-5 promulgated thereunder.

3 47. Defendants, individually and in concert, directly and indirectly, by the use and
4 means of instrumentalities or interstate commerce and/or of the mails, engaged and participated in
5 a continuous course of conduct to conceal adverse material information about the business and
6 future prospects of Ardelyx, as specified herein.

7 48. Defendants employed devices, schemes, and artifices to defraud while in
8 possession of material, adverse nonpublic information and engaged in acts, practices, and a course
9 of conduct, as alleged herein, in an effort to assure investors of Ardelyx's value and performance
10 and continued substantial growth, which included the making of, or participation in the making of,
11 false statements of material facts and omitting to state material facts necessary in order to make
12 the statements made about Ardelyx and its business operations and future prospects, in the light of
13 the circumstances under which they were made, not misleading, as set forth more particularly
14 herein, and engaged in transactions, practices, and a course of business that operated as a fraud
15 and deceit upon the purchasers of Ardelyx securities.

16 49. As described above, Defendants acted with scienter throughout the Class Period in
17 that they either had actual knowledge of the misrepresentations and omissions of material facts set
18 forth herein, or acted with reckless disregard for the truth in that they failed to ascertain and to
19 disclose such facts, even though such facts were available to them. Defendants' material
20 misrepresentations and/or omissions were done knowingly or recklessly and, for the purpose and
21 effect of concealing the Company's results and growth prospects, thereby artificially inflating the
22 price of its securities. As demonstrated by Defendants' omissions and misstatements of the
23 Company's business strategy, Defendants, if they did not have actual knowledge of the
24 misrepresentations and omissions alleged, were reckless in failing to obtain such knowledge by
25 deliberately refraining from taking those steps necessary to discover whether those statements
26 were false or misleading.

1 50. As a result of the dissemination of the materially false and misleading information
 2 and failure to disclose material facts, as set forth above, the market price of Ardelyx securities was
 3 artificially inflated. In ignorance of the fact that market prices of Ardelyx's securities were
 4 artificially inflated, and relying directly or indirectly on the false and misleading statements made
 5 by Defendants, or upon the integrity of the market in which the securities trade, and/or in the
 6 absence of material adverse information that was known to, or recklessly disregarded by,
 7 Defendants, but not disclosed in public statements by Defendants, Plaintiff and the other members
 8 of the Class acquired Ardelyx securities at artificially high prices and were, or will be, damaged
 9 thereby.

10 51. At the time of said misrepresentations and omissions, Plaintiff and the other
 11 members of the Class were ignorant of their falsity and believed them to be true. Had Plaintiff,
 12 the other members of the Class, and the marketplace known the truth regarding the Company's
 13 business, which was not disclosed by Defendants, Plaintiff and the other members of the Class
 14 would not have purchased, or otherwise acquired, their Ardelyx securities, or if they had acquired
 15 such securities, they would not have done so at the artificially inflated prices that they paid.

16 52. By virtue of the foregoing, Defendants have violated §10(b) of the Exchange Act
 17 and Rule 10b-5 promulgated thereunder.

18 53. As a direct and proximate result of Defendants' wrongful conduct, Plaintiff and the
 19 other members of the Class suffered damages in connection with their respective purchases and
 20 sales of the Company's securities.

21 54. This action was filed within two years of discovery of the fraud and within five
 22 years of Plaintiff's purchase of securities giving rise to the cause of action.

23 **For Violations of §20(a) of the Exchange Act**
 24 **(Against the Individual Defendants)**

25 55. Plaintiff repeats and realleges each and every allegation contained above as if fully
 26 set forth herein.

1 56. The Individual Defendants acted as controlling persons of Ardelyx within the
2 meaning of §20(a) of the Exchange Act, 15 U.S.C. §78t(a), as alleged herein. By virtue of their
3 high-level positions, agency, ownership, and contractual rights, and participation in and/or
4 awareness of the Company's operations and/or intimate knowledge of the false financial
5 statements filed by the Company with the SEC and disseminated to the investing public, the
6 Individual Defendants had the power to influence and control, and did influence and control,
7 directly or indirectly, the decision-making of the Company, including the content and
8 dissemination of the various statements that Plaintiff contends are false and misleading. The
9 Individual Defendants were provided with, or had unlimited access to, copies of the Company's
10 reports, press releases, public filings, and other statements alleged by Plaintiff to have been
11 misleading prior to, and/or shortly after, these statements were issued and had the ability to prevent
12 the issuance of the statements or to cause the statements to be corrected.

13 57. In particular, the Individual Defendants had direct and supervisory involvement in
14 the day-to-day operations of the Company and, therefore, are presumed to have had the power to
15 control or influence the particular transactions giving rise to the securities violations, as alleged
16 herein, and exercised the same.

17 58. As set forth above, Ardelyx and the Individual Defendants each violated §10(b) and
18 Rule 10b-5 promulgated thereunder by their acts and omissions, as alleged in this complaint.

19 59. By virtue of their positions as controlling persons, the Individual Defendants are
20 liable pursuant to §20(a) of the Exchange Act. As a direct and proximate result of the Individual
21 Defendants' wrongful conduct, Plaintiff and the other members of the Class have suffered damages
22 in connection with their purchases of the Company's securities.

23 60. This action is filed within two years of discovery of the fraud and within five years
24 of Plaintiff's purchase of securities giving rise to the cause of action.

25 **PRAYER FOR RELIEF**

26 WHEREFORE, Plaintiff prays for relief and judgment, as follows:
27
28

1 A. Determining that this action is a proper class action pursuant to Rule 23(a) and
2 23(b)(3) of the Federal Rules of Civil Procedure on behalf of the Class as defined herein, and a
3 certification of Plaintiff as class representative pursuant to Rule 23 of the Federal Rules of Civil
4 Procedure and appointment of Plaintiff's counsel as Lead Counsel;

5 B. Awarding compensatory and punitive damages in favor of Plaintiff and the other
6 Class members against Defendants, jointly and severally, for all damages sustained as a result of
7 Defendants' wrongdoing, in an amount to be proven at trial, including pre-judgment and post-
8 judgment interest thereon;

9 C. Awarding Plaintiff and other members of the Class their costs and expenses in this
10 litigation, including reasonable attorneys' fees and experts' fees and other costs and disbursements;
11 and

12 D. Awarding Plaintiff and the other Class members such other relief as this Court may
13 deem just and proper.

14 **DEMAND FOR TRIAL BY JURY**

15 Pursuant to Fed. R. Civ. P. 38(b), Plaintiff hereby demands trial by jury of all issues that
16 may be so tried.

17 DATED: July 30, 2021

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